

06020*41210660

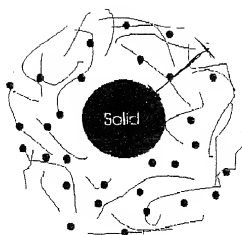


FIGURE 1

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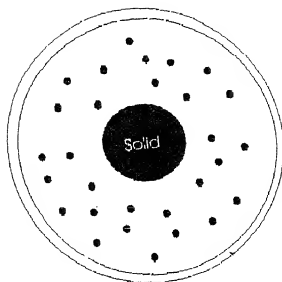


FIGURE 2

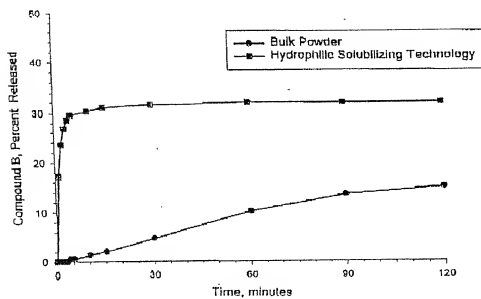


FIGURE 3

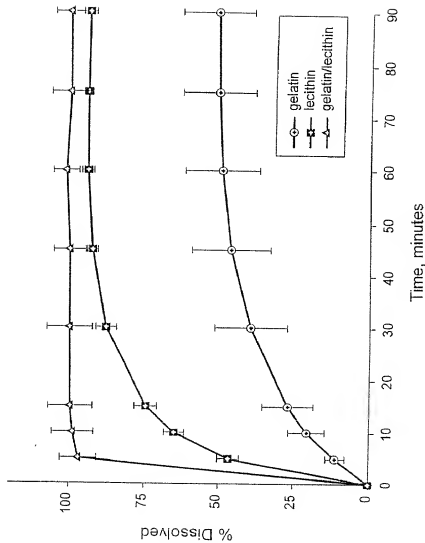


Figure 4: In vitro dissolution of a poorly soluble drug (aqueous solubility = 10 $\mu\text{g/ml}$ at 37°C) formulated with lecithin, gelatin, and the present invention combining lecithin & gelatin.